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In silico improvement of β^3 -peptide inhibitors of p53•hDM2 and p53•hDMX

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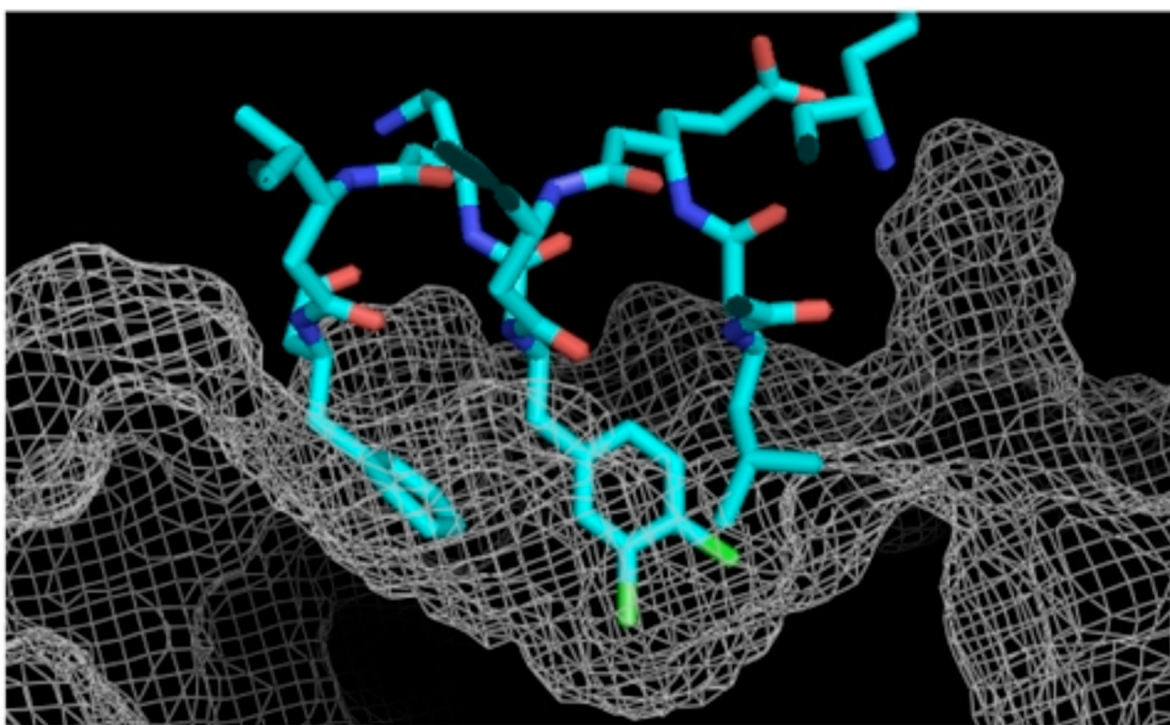
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Abstract

There is great interest in molecules capable of inhibiting the interactions between p53 and its negative regulators hDM2 and hDMX, as these molecules have validated potential against cancers in which one or both oncoproteins are overexpressed. We reported previously that appropriately substituted β^3 -peptides inhibit these interactions and, more recently, that minimally cationic β^3 -peptides are sufficiently cell permeable to upregulate p53-dependent genes in live cells. These observations, coupled with the known stability of β -peptides in a cellular environment, and the recently reported structures of hDM2 and hDMX, motivated us to exploit computational modeling to identify β -peptides with improved potency and/or selectivity. This exercise successfully identified a new β^3 -peptide, **β^3 53-16**, that possesses the highly desirable attribute of high affinity for both hDM2 as well as hDMX and identifies the 3,4-dichlorophenyl moiety as a novel determinant of hDMX affinity.

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Supporting Information Available: Details of computational protocols, CD and binding assays and the complete citation for reference 20. This material is available free of charge via the Internet at <http://pubs.acs.org>



There is great interest in molecules that inhibit interactions between p53 and its negative regulators hDM2 and hDMX, as these molecules have validated potential against cancers that overexpress one or both of these oncoproteins.^{1,2} We reported that substituted β^3 -peptides can inhibit these interactions^{3,4} and, more recently, that minimally cationic β^3 -peptides are sufficiently cell permeable to upregulate p53-dependent genes in live cells.^{5,6} These observations, coupled with the established intracellular stability of β -peptides^{7–9} and the recently reported structures of hDM2¹⁰ and hDMX,¹¹ motivated us to exploit computational methods to identify β -peptides with improved potency and/or selectivity. This exercise successfully identified a new β^3 -peptide, **β 53-16**, that possesses the desirable attribute of high affinity for hDM2 and hDMX and identifies the 3,4-dichlorophenyl moiety as a novel determinant of hDMX affinity.

Our computational modeling began with the application of Visual Molecular Dynamics (VMD)¹² to generate a model of previously reported **β 53-8** bound to the p53 binding site on hDM2 (Figure 1A). In this model, **β 53-8** is bound as a 14-helix that is slightly unwound at the C-terminus, mimicking its conformation in solution.¹³ The three hDM2 hydrophobic pockets occupied in the native structure by the p53 side chains of Leu₂₆, Trp₂₃ and Phe₁₉ are occupied in the modeled complex by the corresponding β^3 -amino acid side chains at positions 3, 6, and 9. An analogous model of **β 53-8** bound to hDMX was also prepared (Figure 1B).¹¹

We then applied a hierarchical computational strategy to search for alternative side chains that would improve packing at one or both interfaces. With the *de novo* design program BOMB¹⁴ we screened over ten thousand **β 53-8** analogs containing substituted aromatic and non-aromatic heterocycles and short hydrocarbon side chains in place of Leu₂₆, Trp₂₃ and Phe₁₉.¹⁰ About 50 candidates were identified by scoring and visualization for evaluation with MCPRO.¹⁵ Binding free energies were predicted *via* Monte Carlo Free Energy Perturbation (MC/FEP) calculations using the OPLS-AA force field¹⁶ for the protein-ligand complex and the TIP4P model for water.¹⁷ In these simulations, the protein backbones remained fixed; the

affinities of the eight most interesting and synthetically accessible compounds (Figure 1C) were subsequently reevaluated in a second round of MC/FEP calculations that permitted backbone motions.¹⁸

The models were first validated by evaluating whether they would predict the large increase in hDM2 affinity realized when the tryptophan side chain at position 6 is replaced by 6-chlorotryptophan (⁶-ClW) (compare **β53-8** and **β53-13**, Figure 1C).¹⁹ The calculations predict that substitution of ⁶-ClW at position 6 should significantly improve binding to hDM2 ($\Delta\Delta G = -2.1 \text{ kcal}\cdot\text{mol}^{-1}$) but not hDMX ($\Delta\Delta G = +1.0 \text{ kcal}\cdot\text{mol}^{-1}$, Figure 2C). These predictions are fully aligned with the experimental results: the stability of the hDM2•**β53-13** complex is significantly higher ($K_d = 30.1 \text{ nM}$, $\Delta G = -10.25 \text{ kcal}\cdot\text{mol}^{-1}$) than that of the hDM2•**β53-8** complex ($K_d = 204 \text{ nM}$, $\Delta G = -9.12 \text{ kcal}\cdot\text{mol}^{-1}$), whereas the stabilities of the analogous hDMX complexes are comparable ($K_d = 1.6$ and $2.1 \text{ }\mu\text{M}$ for **β53-13** and **β53-8**, respectively). The improvement in hDM2 but not hDMX affinity upon substitution of ⁶-ClW is consistent with results observed in the context of previously reported ligands.^{20–23}

The models were further validated by their ability to predict the large increase in hDM2 and hDMX affinity observed for β -peptides containing a central *meta*-trifluoromethyl phenyl substituent (^{CF3}F) when compared with an unsubstituted phenyl ring (compare **β53-12** with **β53-14**, Figure 1C). The calculations predict that the ^{CF3}F side chain should favor binding to both hDM2 and hDMX ($\Delta\Delta G = -4.8$ and $-4.6 \text{ kcal}\cdot\text{mol}^{-1}$, respectively). This increase was also realized experimentally, albeit in an attenuated way: the stability of the hDM2•**β53-12** complex is significantly higher ($K_d = 28.2 \text{ nM}$, $\Delta G = -10.29 \text{ kcal}\cdot\text{mol}^{-1}$) than that of the hDM2•**β53-14** complex ($K_d = 816 \text{ nM}$, $\Delta G = -8.3 \text{ kcal}\cdot\text{mol}^{-1}$); analogous differences are seen for the hDMX complexes (Figure 2).²⁴

Next we examined whether the affinity of **β53-12** could be increased further by substituting the leucine side chain at position 6 with one of eight cyclic and acyclic hydrocarbon alternatives. Although few promising candidates emerged from the BOMB and MC/FEP analyses, we did investigate **β53-17**, in which the Leu side chain is replaced by Ile. This substitution was predicted to slightly favor the binding of both hDM2 and hDMX ($\Delta\Delta G_{\text{bind}} = -0.9$ and $-0.3 \text{ kcal}\cdot\text{mol}^{-1}$, respectively). However, no increase in affinity was observed, and these molecules were not studied further. We note that **β53-17** is significantly less α -helical than **β53-12** as judged by circular dichroism analysis (Figure SI-1). As the computational model does not account for changes in β -peptide secondary structure, it is possible that the observed change in secondary structure accounts for the poor agreement between prediction and experiment in this case. The predictions may be also be affected by uncertainty in the structures of unliganded hDM2 and hDMX as the 23 N-terminal residues of both proteins are only partially resolved due to their flexibility.^{25,26}

Based on these observations, we returned attention to the central side chain of the hDM2/hDMX epitope and evaluated the relative hDM2 and hDMX affinities of hundreds of **β53-12** analogs containing substituted phenylalanine analogs at position 6. This analysis suggested that β -peptides containing either *meta*-chlorophenylalanine or *para*-chlorophenylalanine at this position would show improved affinity for both hDM2 and hDMX when compared with **β53-14** ($-3.5 \text{ kcal}\cdot\text{mol}^{-1} > \Delta\Delta G_{\text{bind}} > -2.5 \text{ kcal}\cdot\text{mol}^{-1}$). Indeed, the stability of the hDM2•**β53-15** complex (*meta*-chloro substituent, Figure 1C) is significantly higher ($K_d = 150 \text{ nM}$, $\Delta G = -9.3 \text{ kcal}\cdot\text{mol}^{-1}$) than that of hDM2•**β53-14**; analogous differences are observed for the hDMX complexes (Figure 2). However, as predicted, the stabilities of the **β53-15** complexes were not greater than those of the **β53-12** complexes. Therefore, since the gains in affinity for the *para*-chlorophenylalanine were predicted to be similar to those of **β53-15**, this additional analog was not tested experimentally.

Finally we examined the effect of a *meta,para*-dichlorophenylalanine side chain at the central position of the recognition epitope (**β53-16**, Figure 1), whose inclusion was predicted to significantly improve affinity for both hDM2 and hDMX compared to **β53-14** ($\Delta\Delta G_{\text{bind}} = -4.4$ and $-5.4 \text{ kcal}\cdot\text{mol}^{-1}$, respectively). Indeed, the stabilities of both hDM2•**β53-16** and hDMX•**β53-16** are significantly higher than the corresponding **β53-14** complexes ($K_d = 27.6 \text{ nM}$ and 155 nM , respectively for **β53-16**, Figure 2). They also equal or exceed the stabilities of the corresponding complexes with **53-12**. Competition fluorescence polarization experiments confirm that **β53-16** competes with p53AD for binding to hDM2 and hDMX and shows improved inhibitory potency towards hDMX (Figure SI-2).

Thus, **β53-16** offers significantly improved affinity for hDMX without loss of affinity for hDM2. Analysis of the MC/FEP simulations suggests more favorable interaction of the dichlorophenyl group with residues 50–54 in hDMX than with equivalent residues 54–58 in hDM2. We subsequently examined whether the affinity of **β53-16** could be improved further upon replacement of the adjacent phenylalanine side chain with one of twelve substituted analogs. This scan failed to identify promising substitutions as the phenylalanine side chain appears to bind tightly to the hydrophobic pocket of both proteins; however minor gains in affinity for both hDM2 and hDMX were predicted for a *para*-fluorophenylalanine substitution ($\Delta\Delta G_{\text{bind}} = -0.4$ and $-0.9 \text{ kcal}\cdot\text{mol}^{-1}$, respectively). No increase in affinity was observed experimentally with this peptide, **β53-18**, so further modification at this position was not pursued (Figure 1, Table 1). **β53-16** represents the highest affinity β^3 -peptide for hDMX reported to date, with significantly higher affinity than the prototypic hDM2 ligand, Nutlin-3. Thus, **β53-16** embodies the pan-specificity of well known peptidic hDM2/hDMX inhibitors^{27,28} without the limitations of protease sensitivity or poor uptake.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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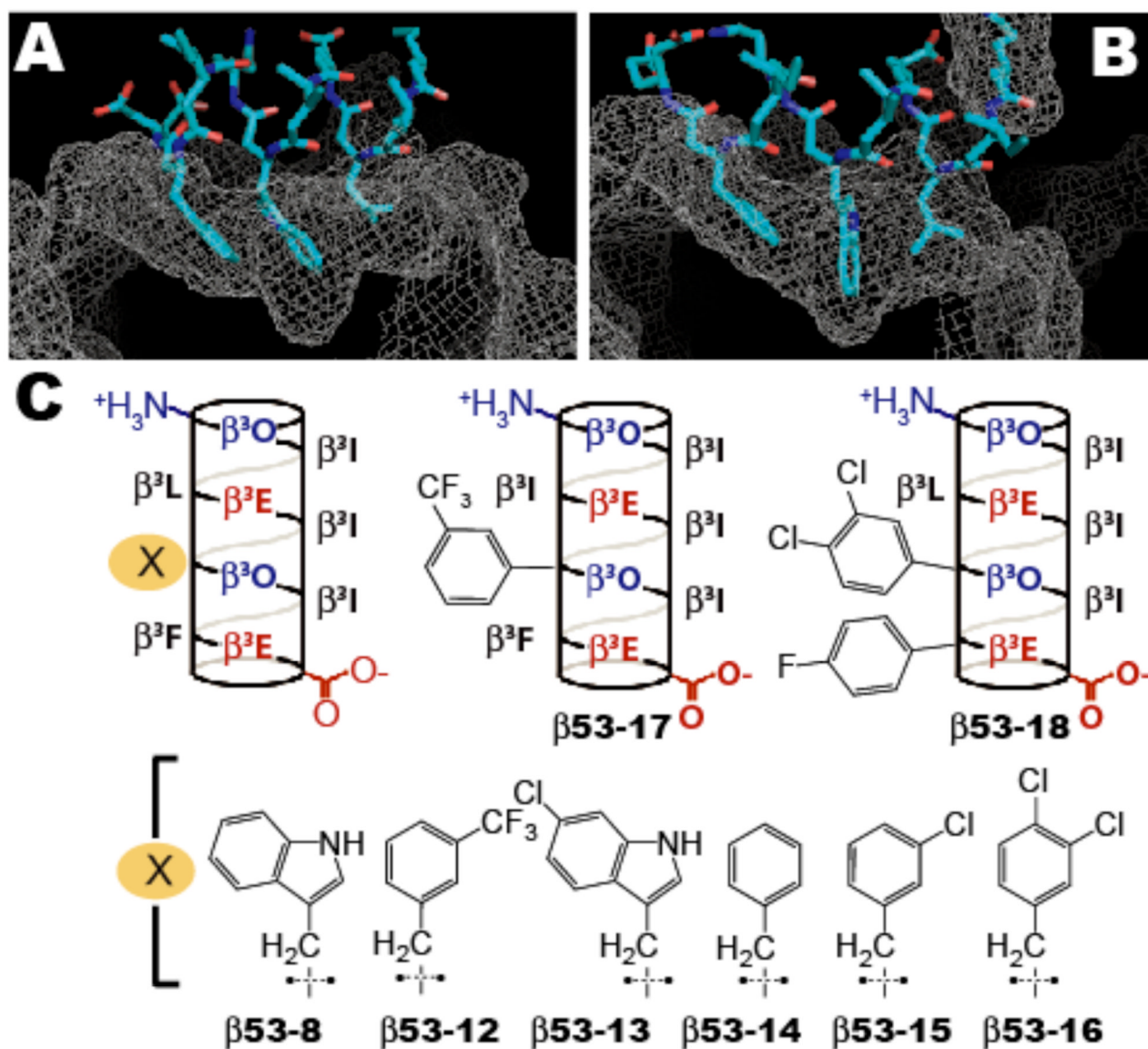
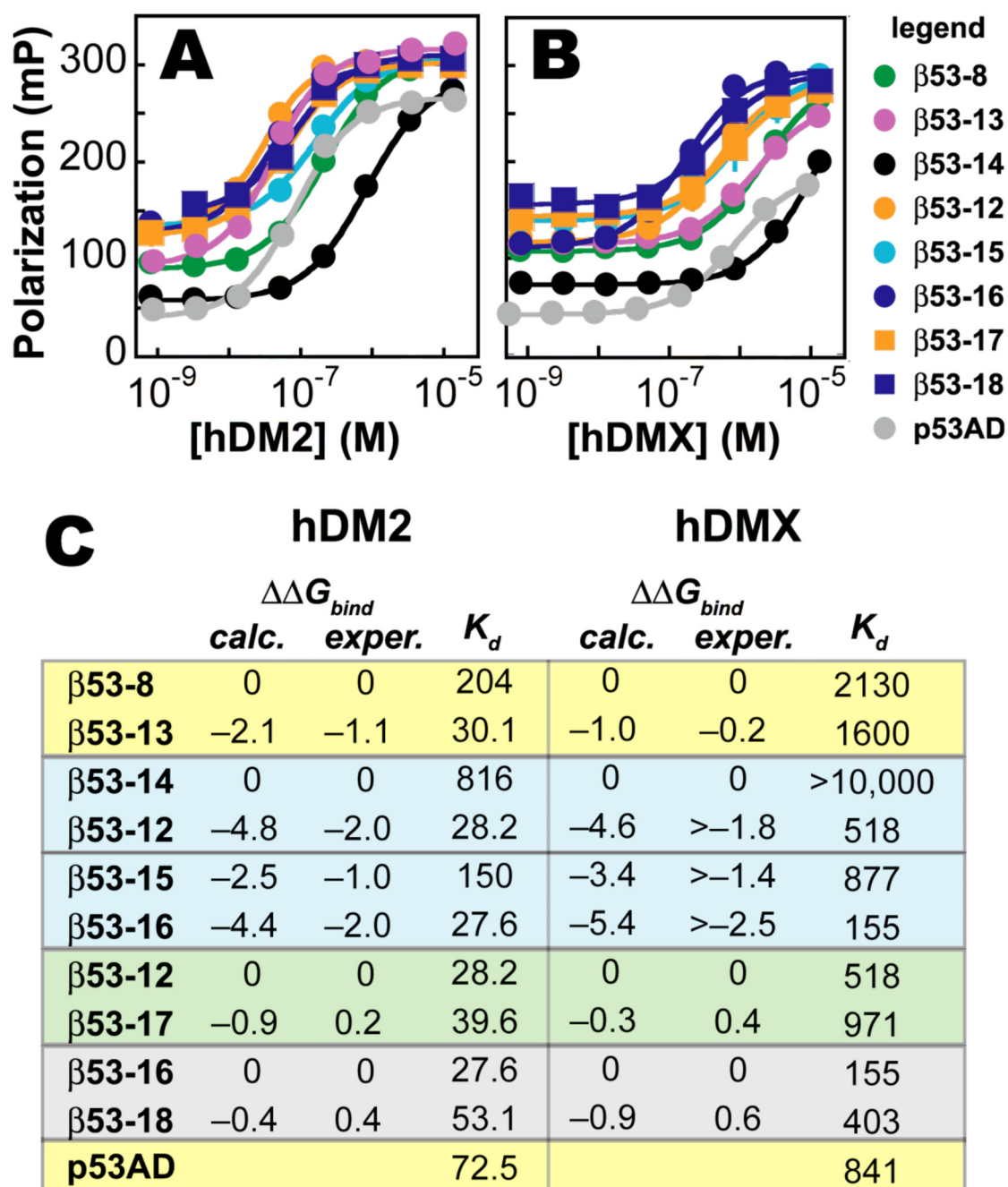


Figure 1.

Computationally generated models of β^{53-8} (blue) in complex with (a) hDM2 and (b) hDMX illustrating differences in binding site topologies. (c) Helical net representations of β^3 -peptides studied herein.

**Figure 2.**

Direct fluorescence polarization analysis of the affinity of each β -peptide shown for (A) hDM2 and (B) hDMX. (C) Comparison of calculated and experimental binding free energies expressed in terms of $\Delta\Delta G_{bind}$ relative to the standard shown ($\text{kcal}\cdot\text{mol}^{-1}$); K_d values in nM units.